# **ZENOL MILD HOT- methyl salicylate poultice Green Cross Corp**

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

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#### **Drug Facts**

methyl salicylate

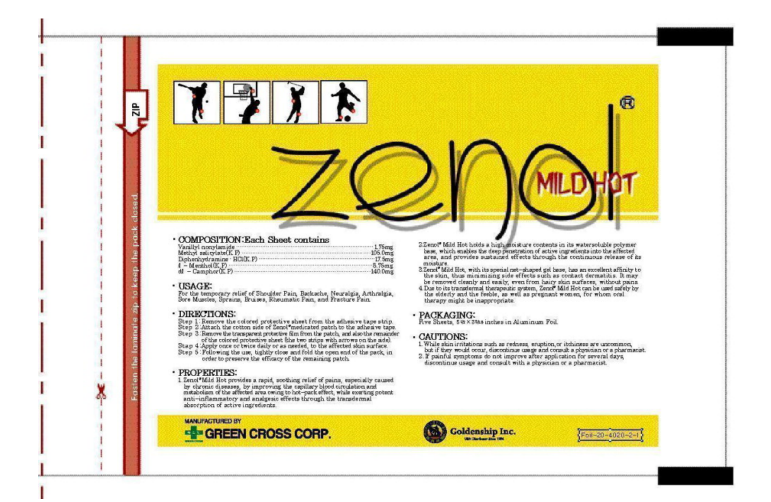
I-menthol, dI-camphor, thymol, diphenhydramine hydrochloride, vanillyl nonylamide, gelatin, concentrated glycerin, zinc oxide, titanium oxide, carboxymethylcellulose sodium, sodium polyacrylate, sodium metaphosphate, purified lanolin, polyoxyetheylene castor oil, phosphoric acid, ferric oxide, rose incense, purified water, nonwoven fabric, polypropylene film

for the temporary relief of shoulder pain, backache, neuralgia etc keep out or reach of the children

step 1: remove the colored protective sheet from the adhesive tape strip step 2: attach the cotton side of Zenol medicated patch to the adhesive tape step 3: remove the transparent protective film from the patch, and also the remainder of the colored protective sheet (the two strips with arrows on the side) step 4: apply one or twice daily or as needed, to the affected skin surface step 5: following the use, tightly close and fold the open end of the pack, in order to preserve the efficacy of the remaining patch

 while skin irritations such as redness, eruption, or itchiness are uncommon, but if they would occur, discontinue usage and consult a physician or a pharmacist
if painful symptoms do no improve after application for several days, discontinue usage and consul with a physician or a pharmacist

for external use only



#### **ZENOL MILD HOT**

methyl salicylate poultice

ı	Product	Information		

Product Type HUMAN OTC DRUG Item Code (Source) NDC:61476-101

Route of Administration TRANSDERMAL

### **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
METHYL SALICYLATE (UNII: LAV5U5022Y) (SALICYLIC ACID - UNII: O414PZ4LPZ)	METHYL SALICYLATE	105 mg in 17.5 g

Inactive Ingredients			
Ingredient Name	Strength		
MENTHOL, UNSPECIFIED FORM (UNII: L7T10EIP3A)			
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET)			
CARBOXYMETHYLCELLULOSE SODIUM, UNSPECIFIED FORM (UNII: K6790BS311)			
GELATIN, UNSPECIFIED (UNII: 2G86QN327L)			
WATER (UNII: 059QF0KO0R)			

Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:61476-101- 01	17.5 g in 1 PATCH; Type 0: Not a Combination Product	12/03/2013	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph not final	part348	12/03/2013	02/02/2023	

## Labeler - Green Cross Corp (687760561)

Establishment				
Name	Address	ID/FEI	Business Operations	
Green Cross Corp		689852033	manufacture(61476-101)	

Revised: 11/2021 Green Cross Corp